

UNITED STATES ENVIRONMENTAL PROTECTION AGENCY
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FILE

OPP OFFICIAL RECORD
HEALTH EFFECTS DIVISION
SCIENTIFIC DATA FILES
EPA-600/3-98-011OFFICE OF
PREVENTION, PESTICIDES, AND
TOXIC SUBSTANCES**MEMORANDUM**

DATE: May 4, 1998

SUBJECT: ID#97CA0036. SECTION 18 EXEMPTION FOR THE USE OF
**MYCLOBUTANIL ON PEPPERS (BELL and NON - BELL) IN NEW
MEXICO.**

DP Barcode:	D244822	Caswell#:	723K
PRAT Case#:	289836	Chemical#:	128857
Trade Name:	NOVA® 40W	40 CFR:	§180.443
EPA Reg#:	707-221	Class:	Fungicide

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MUIERB/RD (7505C)**INTRODUCTION**

The New Mexico Department of Agriculture has proposed a specific exemption for the use of myclobutanil on peppers (bell and non-bell) for control of powdery mildew (*Oidiopsis taurica*). This is the third §18 request for this use. The proposed program will entail application of 15,000 pounds of Nova® 40W (12,000 lbs ai) on 30,000 acres statewide from June 1, 1998 until October 15, 1998.

SUMMARY

The following restriction should be added to the label for the requested Section 18 on peppers: Myclobutanil treated fields can be rotated at any time to crops which are included on a myclobutanil label. All other crops may be planted one year following application of myclobutanil.

Occupational exposure and aggregate risk estimates do not exceed HED's level of concern (provided the following restrictions are put on the label: 1) The type of flagger utilized, during aerial application, must be indicated on the label (e.g. whether it is mechanical or a worker), if the flagger is not mechanical, then the flagger must wear long pants and a long-sleeved shirt with shoes and socks; and 2) to conduct residential applications on different days, e.g. lawn application done on a different day than flower or tree applications).

Therefore, provided the above statement regarding rotational crops is added to the Section 18 label, HED has no objection to the issuance of this Section 18 exemption for the use of myclobutanil on peppers (bell and non-bell) in the State of New Mexico. The following time-limited tolerances for the combined residues of myclobutanil [α -butyl- α -(4-chlorophenyl)-1*H*-1,2,4-triazole-1-propanenitrile] plus its alcohol metabolite [α -(3-hydroxybutyl)- α -(4-chlorophenyl)-1*H*-1,2,4-triazole-1-propanenitrile] (free and bound) should be established to support this Section 18 exemption:

peppers (bell and non-bell) 1.0 ppm

TOXICOLOGICAL ENDPOINTS

DIETARY

1. *Acute Toxicity*

None. For acute dietary risk assessment, the Hazard ID Assessment Review Committee (HIARC) did not recommend an acute dietary endpoint.

2. *Chronic Toxicity*

RfD = 0.025 mg/kg/day. The RfD is currently established to be 0.025 mg/kg/day based on the NOEL from the chronic feeding study in the rat (2.49 mg/kg/day; MRID #00165247) and a safety factor of 100 [10 for intraspecies and 10 for interspecies]. The LOEL for the chronic rat feeding study is 9.84 mg/kg/day based on decreased testicular weight and increased testicular atrophy. The HIARC noted that the dose of 2.49 mg/kg/day established in the above study is supported by the Parental Systemic Toxicity NOEL and LOEL established in the Two-Generation reproduction study in rats. In that study the NOEL was 2.5 mg/kg/day and the LOEL was 10 mg/kg/day. The Committee determined that the 10 x factor to account for enhanced sensitivity of infants and children (as required by FQPA) should be removed. A UF of 100 is adequate because of the following:

- (i) Developmental toxicity studies showed no increased sensitivity in fetuses as compared to maternal animals following *in utero* exposures in rats and rabbits.
- (ii) A two generation reproduction toxicity study in rats showed no increased sensitivity in pups that were compared to adults.
- (iii) The toxicology data base is complete and there are no data gaps.

This decision was confirmed by the *ad hoc* FQPA Safety Factor Committee (R. Keigwin and W. Burnam, personal communication). The Joint Meeting on Pesticide Residues (JMPR) established an ADI (RfD) of 0.03 mg/kg/day.

NON-DIETARY

1. *Short-Term Toxicity*

For short-term Margin of Exposure (MOE) calculations, the HIARC recommended use of the systemic NOEL of 100 mg/kg/day [HDT] from the 28-day dermal toxicity study in rats (MRID# 266080). There was no LEL in the study.

2. *Intermediate-Term Toxicity*

For intermediate-term MOE calculations, the HIARC recommended use of the reproductive NOEL of 10 mg/kg/day based on atrophy of the testes and prostate as well as an increase in the number of stillborns and a decrease in pup weight gain during lactation at the LOEL of 50 mg/kg/day (LOEL) from the 2-generation reproduction study in rats (MRID# 00143766, 00149581).

3. *Chronic Toxicity*

The HIARC determined that a chronic toxicity endpoint and risk assessment for myclobutanil is not required for workers.

4. *Dermal Penetration*

For short-term MOE calculations, a dermal toxicity study was used, so dermal penetration data were not required. The HIARC determined that a dermal absorption factor of 100% should be used for risk assessment because 1) a dermal absorption study was not available with the technical and 2) a dermal absorption factor could not be estimated due to the lack of comparative NOELs/LOELs from oral and dermal toxicity studies in the same species with the technical. The dermal absorption factor is required for Intermediate and Long-Term dermal risk assessment since oral doses were selected for these exposure periods. Dermal absorption is not required for Short-Term dermal exposure risk assessment since a dermal dose from a 28-day dermal toxicity study was selected for this time period.

CANCER

Myclobutanil is classified as Category E: not carcinogenic in two acceptable animal studies. Q_1^* is not applicable.

EXPOSURES AND RISKS

In examining aggregate exposure, FQPA directs EPA to consider available information concerning exposures from the pesticide residues in food and all other non-occupational exposures. The primary non-food sources of exposure the Agency looks at include drinking

water (whether from groundwater or surface water), and exposure through pesticide use in gardens, lawns, or buildings (residential and other outdoor and indoor uses). In evaluating food exposures, EPA takes into account varying consumption patterns of major identifiable subgroups of consumers, including infants and children.

1. From Food and Feed Uses:

Tolerances have been established (40 CFR 180.443) for the residues of myclobutanil [alpha-butyl-alpha-(4-chlorophenyl)-1H-1,2,4-triazole-1-propanenitrile] and its metabolite alpha-(3-hydroxybutyl)-alpha-(4-chlorophenyl)-1H-1,2,4-triazole-1-propanenitrile (free and bound), expressed as myclobutanil, in or on a variety of raw agricultural commodities and processed commodities at levels ranging from 0.02 ppm in cottonseed to 25.0 ppm in raisin waste. Meat, milk, poultry and egg tolerances have been established at levels ranging from 0.02 ppm to 1.0 ppm.

Acute Risk. The HIARC did not recommend an acute dietary toxicological endpoint so an acute dietary risk assessment is not required (10/21/97 meeting).

Chronic Risk. In conducting this chronic dietary (food only) risk assessment, HED has made somewhat conservative assumptions. With the exceptions of bananas for which a level representing residues in pulp rather than the whole banana was used and selected commodities which were corrected for percent crop treated, all commodities having myclobutanil tolerances will contain myclobutanil and metabolite residues and those residues will be at the level of the established tolerance. This results in an overestimate of human dietary exposure. For bananas, the level of 0.8 ppm was used in the dietary risk assessment rather than the proposed tolerance of 4.0 ppm on bananas since residues in the pulp will not exceed 0.8 ppm. Percent crop-treated estimates were utilized for selected commodities included in the assessment. Thus, in making a safety determination for this tolerance, EPA is taking into account this partially refined exposure assessment.

The existing myclobutanil tolerances (published, pending, and including the necessary Section 18 tolerances) for crops other than bananas and the anticipated residues on bananas result in an Anticipated Residue Contribution (ARC) that is equivalent to the following percentages of the RfD:

Population Subgroup	ARC _{food} (mg/kg/day)	%RfD
U.S. Population (48 states)	0.004274	17%
Nursing Infants (<1 year old)	0.006359	25%
Non-Nursing Infants (<1 year old)	0.018836	75%
Children (1-6 years old)	0.011492	46%
Children (7-12 years old)	0.006910	28%
Northeast Region	0.004566	18%
Western Region	0.004870	19%
Hispanics	0.005065	20%
Non-Hispanic Others	0.004441	18%

The subgroups listed above are: (1) the U.S. population (48 states); (2) those for infants and children; and, (3) the other subgroups for which the percentage of the RfD occupied is greater than that occupied by the subgroup U.S. population (48 states).

2. *From Drinking Water:*

Based on information in the EFED One Liner Database (updated: 12/20/94), myclobutanil is persistent and not considered mobile in soils with the exception of sandy soils. Data are not available for its metabolite alpha-(3-hydroxybutyl)-alpha-(4-chlorophenyl)-1H-1,2,4-triazole-1-propanenitrile. There is no established Maximum Contaminant Level for residues of myclobutanil in drinking water (Safe Drinking Water Hotline - personal communication 5/14/97). No Health Advisory Levels for myclobutanil in drinking water have been established. The "Pesticides in Groundwater Database" (EPA 734-12-92-001, September 1992) has no information concerning myclobutanil.

The Environmental Fate and Effects Division (D239591, Douglas Urban, 11/4/97) has provided estimates of ground and surface water concentrations for myclobutanil based on the label rate of 0.65 lbs a.i./acre and assuming 15 applications per season. (The water numbers were based on turf.) The surface water numbers are based on the results of GENEEC model run. The ground water numbers are based on a screening tool, SCI-GROW, which tends to overestimate the true concentrations in the environment.

Surface water EEC [based on the results of a GENEEC (Version 1.2, 5/3/95) model run]

Acute = 145.96 ppb (0.14596 ppm or mg/L)(maximum initial concentration)

Chronic = 118.6 ppb (0.1186 ppm or mg/L)(average 56-day concentration)

NOTE: OPP policy allows the 90/56-day GENEEC value to be divided by 3 to obtain a value for chronic risk assessment calculations. Therefore, the surface water value for use in the chronic risk assessment would be 0.04 ppm or mg/L.

Ground water EEC (SCI-GROW, Lotus 1-2-3 spreadsheet)

3.6 ppb (0.0036 ppm or mg/L) (use for both acute and chronic)

Chronic exposure from surface water is calculated below. Chronic exposure from ground water is lower.

OPP has calculated drinking water levels of concern (DWLOCs) for chronic (non-cancer) exposure to be **0.7 ppm** for the U.S. population, **0.6 ppm** for Hispanics, and **0.06 ppm** for non-nursing infants (<1 year old). To calculate the DWLOC for chronic (non-cancer) exposure relative to a chronic toxicity endpoint, the chronic dietary food exposure (from DRES) was subtracted from the RfD to obtain the acceptable chronic (non-cancer) exposure to myclobutanil in drinking water.

Note: The following formula was used to convert maximum allowable water exposure to ppb. DWLOCs were then calculated using default body weights (70 kg - adult, 10 kg - child) and drinking water consumption figures (2 L - adult, 1 L child).

$$DWLOC (\mu\text{g/L}) = \frac{\text{water exposure (mg/kg/day)} \times (\text{body weight})}{\text{consumption (L)} \times 10^{-3} \text{ mg}/\mu\text{g}}$$

The estimated average concentration of myclobutanil in surface water is **0.04 ppm**. Chronic concentrations in ground water are not expected to be higher than the acute concentrations. The estimated average concentrations of myclobutanil in surface water are less than OPP's levels of concern for myclobutanil in drinking water as a contribution to chronic aggregate exposure. Therefore, taking into account the present uses and uses proposed in this action, OPP concludes with reasonable certainty that residues of myclobutanil in drinking water (when considered along with other sources of exposure for which OPP has reliable data) would not result in unacceptable levels of aggregate human health risk at this time.

OPP bases this determination on a comparison of estimated concentrations of myclobutanil in surface waters and ground waters to back-calculated "levels of concern" for myclobutanil in drinking water. These levels of concern in drinking water were determined after OPP has considered all other non-occupational human exposures for which it has reliable data, including all current uses, and uses considered in this action. The estimates of myclobutanil in surface waters are derived from water quality models that use conservative assumptions (health-protective) regarding the pesticide transport from the point of application to surface and ground water. Because OPP considers the aggregate risk resulting from multiple exposure pathways associated with a pesticide's uses, levels of concern in drinking water may vary as those uses change. If new uses are added in the future, OPP will reassess the potential impacts of myclobutanil on drinking water as a part of the aggregate risk assessment process.

3. *From Non-Dietary Uses:*

Myclobutanil is currently registered for outdoor residential and greenhouse use on annuals and perennials, turf, shrubs, trees, and flowers (Reference Files System/OPP LAN, date searched: 6/5/97). HED has determined that these uses do not constitute a chronic exposure scenario, but may constitute a short- to intermediate-term exposure scenario (**Note: the intermediate-term potential exposure would come from Post-application (dermal for adult; and dermal + ingestion of soil only, due to the persistence of the pesticide in soil, for toddlers)**). Other intermediate-term exposure scenarios are unlikely as dissipation is strongly influenced by the growth of the grass which needs weekly mowing (more frequently in spring) and most dissipation studies on lawns show considerable tailing off of residues by day 3 or 4; thus, the expectation of significant residues is very unlikely.

Homeowner-use Products

End-use products containing the active ingredient, myclobutanil, are marketed for homeowner use. The homeowner use with the greatest potential for exposure takes the form of small scale lawn application (**other additional application uses are on roses, flowers, ornamental shrubs and trees**) of a soluble concentrate with a hose-end, backpack, or trigger bottle sprayer. Application of these products is recommended at two week intervals. Short-term (and not intermediate-term exposures, because of the amount of time it takes to mix, load, and apply this

product) exposure is considered only. Short-term exposure, pre- and during application, will be considered an aggregate potential exposure: a summation of this exposure will include exposure levels for: the mixer + loader + applicator + Post-application on day zero (day of application). Short- and intermediate-term exposure will be considered during post-application (*Note:* Intermediate-term exposure is addressed only during post-application scenarios).

Handler Exposures and Assumptions

HED has determined that there is potential for exposures to applicators and handlers during usual homeowner use-patterns associated with myclobutanil. Based on the use patterns, three exposure scenarios with the greatest potential for exposure are considered: 1) loading and application of a soluble concentrate product by low pressure handwand sprayer (trigger bottle sprayer); 2) loading and application of a soluble concentrate product by backpack; and 3) loading and application of a soluble concentrate product by garden hose end sprayer.

Short-term dermal exposure assessments using the Pesticide Handlers Exposure Database (PHED) Version 1.1 surrogate data and baseline risk calculations for homeowners are presented in Table 1. Table 2 summarizes the caveats (e.g., data confidence) and parameters specific to each exposure scenario and corresponding risk assessment.

TABLE 1. Baseline Short-Term Exposure and Risk Assessments for Homeowner Use of Myclobutanil

Exposure Scenario	Baseline Dermal + Inhalation Unit Exposure (mg/lb ai) ^a	Maximum Application Rate (lb ai/acre) ^b	Maximum Acres/Day ^c	Total Daily Exposure (mg ai/day) ^d	Total Daily Dose (mg ai/kg/day) ^e	Short-Term MOE ^f
					BW = 60 kg	
1. Load/Apply Soluble Concentrate Using Low Pressure Handwand	100.03 100.03 (0.03) ^h	0.63 1.7 ^a 0.63	0.50 0.50 0.50	32 85 ^a (9.4X10 ⁻³) ^h	1 1.9 ^a (1.6X10 ⁻⁴) ^h	100 53 ^a (62000) ⁱ
2. Load/Apply Soluble Concentrate Using Backpack	5.1 (0.03) ^h	0.63 0.63	0.50 0.50	1.6 (9.4X10 ⁻³) ^h	0.49 (1.6X10 ⁻⁴) ^h	200 (62000) ⁱ
3. Load/Apply Soluble Concentrate Using Garden Hose End Sprayer	30.01 (0.01) ^h	0.63 0.63	0.50 0.50	9.4 (3.2X10 ⁻³) ^h	0.62 (5.3X10 ⁻⁵) ^h	160 (190000) ⁱ

a Baseline unit exposure (dermal + inhalation), taken from PHED Version 1.1 data in the Draft Standard Operating Procedures (SOPs) for Residential Exposure Assessments dated December 18, 1997, represents short pants, short sleeve shirt, no gloves, and open loading.

Note: that for some PHED data, correction factors were applied to arrive at the baseline scenario.

b Application rate comes from maximum rates found on the Myclobutanil labels.

c Daily acres treated values are from Draft Standard Operating Procedures (SOPs) for Residential Exposure Assessments dated December 18, 1997, estimates of acreage that could be treated in a single day for each exposure scenario of concern.

d Total Daily Exposure (mg ai/day) = Unit exposure (mg/lb ai) x Application Rate (lbs ai/acre) x Acres Treated.

e Total Daily Dose (mg/kg/day) = Daily (dermal+inhalation) Exposure (mg a.i./day) + Post-application exposure on day of application (= 28 mg a.i./day) /body weight (BW kg). See calculation for Post-application exposure below.

f Margin of Exposure (MOE) = NOEL (mg/kg/day)/Daily Dose (mg/kg/day); NOEL = 100mg/kg/day; MOE for 60kg

g Worst case day (based on label) of potential exposures = mixer/loader and application of lawn +roses+tree+flowers.

h Inhalation unit exposure only, taken from PHED Version 1.1 data in the Draft Standard Operating Procedures (SOPs) for

Residential Exposure Assessments dated December 18, 1997, represents short pants, short sleeve shirt, no gloves, and open loading. For Total Daily Dose (mg/kg/day) = Daily(inhalation) Exposure (mg a.i./day)/body weight (BW kg).

Short-term, inhalation exposure only, MOE based on a NOEL = 10 mg/kg/day.

Table 2. Exposure Scenario Descriptions for Selected Residential Uses of Myclobutanil

Exposure Scenario (Number)	Data Source	Standard Assumption ^a	Comments
Mixer/Loader/Applicator Descriptors			
Load/Apply Soluble Concentrate Using Low Pressure Handwand (1)	PHED V1.1	0.50 Acres	Baseline: Low confidence (9-80 replicates of ABC grade data) for dermal exposure. Medium confidence (80 replicates of ABC grade data) for inhalation.
Load/Apply Soluble Concentrate Using Backpack (2)	PHED V1.1	0.50 Acres	Baseline: Low confidence (9-11 replicates of AB grade data) for dermal exposure. Low confidence (11 replicates of A grade data) for inhalation.
Load/Apply Soluble Concentrate Using Garden Hose End Sprayer (3)	PHED V1.1	0.50 Acres	Baseline: Low confidence (8 replicates of C and E grade data) for dermal exposure. Low confidence (8 replicates of C grade data) for inhalation. <i>Based on one study.</i>

^a Standard Assumptions based on Draft Standard Operating Procedures (SOPs) for Residential Exposure Assessments dated December 18, 1997. Baseline dermal exposure is based on the worker wearing short pants, short sleeve shirt, and no gloves.

Formulas for determining daily (dermal+Inhalation)exposure and risk to handlers are as follows:

$$\text{Daily Exposure (mg a.i./day)} =$$

$$\text{Unit Exposure (mg/lb)} \times \text{Use Rate (lb a.i./acre)} \times \text{Maximum Area Treated (acres/day)}$$

$$\text{Daily (Dermal + Inhalation; or Inhalation only) Dose (mg a.i./kg bw/day)} =$$

$$\frac{\text{Daily Exposure (mg a.i./day)}}{\text{Body Weight (kg)}}$$

$$\text{Margin of Exposure (MOE)} =$$

$$\frac{\text{NOEL (mg/kg/day)}}{\text{Daily (Dermal + Inhalation; or Inhalation only) Dose (mg/kg/day)}}$$

The following are important assumptions used in the residential exposure assessments:

- For the short-term exposure assume exposed person's body weight is 60 kg; For the toddler (age 3) assume exposed body weight is 15 kg;
- Footnotes for Table 1 include other assumptions.

Homeowner Post-Application Exposures and Assumptions

The potential for post-application homeowner exposure exists. For example, potential exposures would be expected following applications to lawns and ornamental garden sites. There are no chemical-specific data to use in assessing these potential exposures; therefore, a range finder post-application exposure and risk assessment was performed (Table 3). The assessment uses typical transfer coefficients (Tc); for Adults = 10,000 cm²/hr (high activity for 4 hrs, Tier II.) and for Toddlers = 8,700 cm²/hr (for 2hrs. default tier I). It also utilizes dislodgeable foliar residues (DFR) derived from the application rate and an estimated 10 percent [less conservative than the default 20 percent (which is based on foliar wash), but still much more conservative than the California's roller method study of rate available as dislodgeable (which had an average of 1-2 % of rate available as DFR)]. EPA believes that exposures following soluble concentrate applications with a low pressure handwand to plants, such as lawn-turfgrass, are likely to represent a reasonably conservative post-application exposure estimate to homeowners and children. Total aggregate short-term exposure was calculated for adults and toddlers (for toddlers include dermal + incidental non-dietary ingestion (hand to mouth; surface area for one hand=175cm²)). Isolated scenarios for short-term exposure for toddlers also include; ingestion of treated turfgrass, and ingestion of treated soil. Intermediate-term exposure for adults will be a mean value of a 14-day exposure scenario. For intermediate-term, total aggregate exposure for toddlers will include only, a mean value of a 14-day exposure scenario + the ingestion of soil. **Chemical-specific dissipation data and residential use/usage information are required to further refine these post-application exposure estimates.**

Table 3. Surrogate Post-application Range-Finder Assessment.

DAT ^a	DFR (µg/cm2) ^b	Dermal Dose (mg/kg/day) ^c		Adult Short- Term MOE ^d	Adult Intermediate- Term MOE ^d	Toddler Short- Term MOE ^d	Toddler Intermediate- Term MOE ^d
		BW = 60 kg	BW = 15 kg				
Exposure Activities (Tc = 10,000 cm ² /hr (adults-tierII) ; For toddlers 8,700 (default) cm ² /hr) ^e							
0	0.71	0.47 ^c ₁	0.85 ^c ₂	210 ^d ₁	N/A	120 ^d ₂	N/A
0	0.71	N/A	1.2 X10 ^{-3c} ₃	N/A	N/A	83,000 ^d ₃	N/A
0	N/A	N/A	3.2 X10 ^{-5c} ₄	N/A	N/A	3.1 X10 ^{6d} ₄	N/A
0-14	0.71	0.04	N/A	N/A	250 ^d ₅	N/A	N/A
0-14	0.71	N/A	6.1 X10 ⁻²	N/A	N/A	N/A	160 ^d ₆

a DAT is days after treatment based on an application rate of 1.44X10⁻⁵ lb ai/ ft².

b DFR (µg/cm²) = Rate (lb ai/ft²) x (weight conversion factor to convert the lbs a.i. in the application rate to µg for the DFR value = 4.54x 10⁸ µg/lb) x (area unit conversion factor = 1.08 x 10⁻³ ft²/cm²) x percent (10 percent assumed) of rate available as dislodgeable

c **Dermal Dose** (mg/kg/day): c₁= For Adult Females, = **Dermal Dose** (mg/kg/day) = [DFR (µg/cm²) x Tc (cm²/hr) x (1 mg/1,000 µg unit conversion) x 4 hours/day(tier II)] / Body Weight (BW kg); c₂= For Toddlers, = **Dermal Dose** (mg/kg/day) = [DFR (µg/cm²) x Tc (cm²/hr) x (1 mg/1,000 µg unit conversion) x 2 hours/day(default) / Body Weight (BW kg) + **Incidental non-dietary ingestion of pesticide residues on residential lawns from hand to mouth transfer**=DFR X surface area of one hand (175cm²/event) X frequency of hand to mouth activity(1.56events/hr) X exposure time (2hrs/day) X weight unit conversion factor (1mg/1000 µg) /BW; **Isolated incidents for Toddlers, 1)** c₃= **Ingestion of treated turfgrass** = grass (and plant matter) residue on day of application (µg/cm²) X ingestion rate of grass (25cm²/day) X conversion factor (mg/1000 µg)

- /BW; and 2) c_d =, **Ingestion of treated soil** =soil residue on day of application ($\mu\text{g/g}$) X ingestion rate of soil (100mg/day) X weight unit conversion factor($1\text{g}/1\times 10^6 \mu\text{g}$) /BW
- d MOE = NOEL (mg/kg/day)/Dermal Dose (mg/kg/day); Short-term NOEL=100mg/kg/day and Intermediate-term NOEL=10mg/kg/day. d_1 =Short-term, Adult Females ; d_2 = For Toddlers, Short-term exposure, is an aggregate exposure scenario, which includes dermal+ incidental non-dietary ingestion hand to mouth. Other short-term exposure for toddlers, isolated incidents; d_3 = 1) ingestion of treated turf, & d_4 = 2) ingestion of treated soil. **Intermediate-term exposure:** for the adult, d_5 = a mean value (10% DFR= 0.71) based on 14 days with a 10% decrease each day after the day of application (day 0).; **for toddlers**, d_6 = a mean value (10% DFR) based on 14 days with a 10% decrease each day after day 0 + ingestion of treated soil (application rate on Day 0). **Note:** Ingestion of treated soil times 6 months of application exposure, this is a very conservative estimate due to loss of pesticide through rain fall dilution, soil erosion, etc.. **Exposure from treated soil only** = $3.15\times 10^{-5}\text{mg/kg/day} \times 2 \text{ applications/month} \times 6 \text{ months} = 3.8\times 10^{-4}\text{mg/kg/day}$; **MOE** = $10\text{mg/kg/day}/3.8\times 10^{-4}\text{mg/kg/day} = 26,315$.
- e The upper percentile dermal transfer coefficient is assumed to be 43,000 cm^2/hr for adults (default, tier I) and 8,700 cm^2/hr for toddlers (default, tier I).

4. *From Cumulative Exposure To Substances with a Common Mechanism of Toxicity*

Myclobutanil is a member of the triazole class of systemic fungicides (*The Pesticide Book*, 4th ed., 1994). Other triazoles include bitertanol, cyproconazole, diclobutrazole, difenoconazole, diniconazole, fenbuconazole, flusilazole, hexaconazole, penconazole, propiconazole, tebuconazole, tetraconazole, triadimefon, and triadimenol.

Section 408(b)(2)(D)(v) of the Food Quality Protection Act requires that, when considering whether to establish, modify, or revoke a tolerance, the Agency consider "available information" concerning the cumulative effects of a particular pesticide's residues and "other substances that have a common mechanism of toxicity." The Agency believes that "available information" in this context might include not only toxicity, chemistry, and exposure data, but also scientific policies and methodologies for understanding common mechanisms of toxicity and conducting cumulative risk assessments. For most pesticides, although the Agency has some information in its files that may turn out to be helpful in eventually determining whether a pesticide shares a common mechanism of toxicity with any other substances, EPA does not at this time have the methodologies to resolve the complex scientific issues concerning common mechanism of toxicity in a meaningful way. EPA has begun a pilot process to study this issue further through the examination of particular classes of pesticides. The Agency hopes that the results of this pilot process will increase the Agency's scientific understanding of this question such that EPA will be able to develop and apply scientific principles for better determining which chemicals have a common mechanism of toxicity and evaluating the cumulative effects of such chemicals. The Agency anticipates, however, that even as its understanding of the science of common mechanisms increases, decisions on specific classes of chemicals will be heavily dependent on chemical specific data, much of which may not be presently available.

Although at present the Agency does not know how to apply the information in its files concerning common mechanism issues to most risk assessments, there are pesticides as to which the common mechanism issues can be resolved. These pesticides include pesticides that are toxicologically dissimilar to existing chemical substances (in which case the Agency can conclude that it is unlikely that a pesticide shares a common mechanism of activity with other substances) and pesticides that produce a common toxic metabolite (in which case common mechanism of activity will be assumed).

EPA does not have, at this time, available data to determine whether myclobutanil has a common

mechanism of toxicity with other substances or how to include this pesticide in a cumulative risk assessment. For the purposes of these tolerance actions, therefore, EPA has not assumed that myclobutanil has a common mechanism of toxicity with other substances.

DETERMINATION OF SAFETY FOR U.S. POPULATION

1. *Acute Aggregate Risk*

This risk assessment is not required as the HIARC did not identify any acute dietary risk endpoints.

2. *Chronic Aggregate Risk*

Chronic Aggregate Exposure and Risk. Using the partially refined exposure assumptions described above, HED has concluded that aggregate exposure (food, water, and residential) to myclobutanil will not exceed HED's level of concern. For the U.S. population, 17% of the RfD is occupied by dietary (food) exposure. The estimated average concentrations of myclobutanil in surface and ground water are less than OPP's levels of concern for myclobutanil in drinking water as a contribution to chronic aggregate exposure. Therefore, OPP concludes with reasonable certainty that residues of myclobutanil in drinking water do not contribute significantly to the aggregate chronic human health risk at the present time considering the present uses and uses proposed in this action. HED has determined that the outdoor registered uses of myclobutanil would not fall under a chronic exposure scenario. HED concludes that there is a reasonable certainty that no harm will result from aggregate chronic exposure to myclobutanil residues.

3. *Short-Term Aggregate Risk*

The short-term NOEL for dermal exposure is based on a dermal exposure toxicity study. Since the NOEL is based on a dermal study, oral exposures generally cannot be used directly to calculate a short-term aggregate exposure. However, as the HIARC determined that a dermal absorption factor of 100% should be used for risk assessment, oral exposures need not be multiplied by a modifying factor (converted to dermal equivalents) so that they can be compared to the dermal endpoint.

The chronic dietary exposure and calculated dietary MOE is shown below for the U.S. Population.

Subgroup	ARC (from DRES) (mg/kg/day)	Calculated Dietary MOE (from DRES)
U.S. Population (48 states)	0.004274	24000

Calculations:

$$\text{Dietary MOE} = \frac{\text{Short-term NOEL}}{\text{Chronic dietary exposure}}$$

$$= \frac{100 \text{ mg/kg/day}}{0.004274 \text{ mg/kg/day}} = 24,000$$

The dermal residential exposure for different scenarios and aggregate short-term MOEs is shown below for the U.S. Population (48 states).

Exposure scenario	Calculated Dietary MOE (from DRES)	Total Residential Exposure (from Table 1) (mg/kg/day)	Total Residential MOE	Total Short-term MOE (Dietary + Residential)
Load/Apply Soluble Concentrate Using Low Pressure Handwand	24,000	1	100	100
	24,000	1.9 ^a	53 ^a	53 ^a
Load/Apply Soluble Concentrate Using Backpack	24,000	0.49	200	200
Load/Apply Soluble Concentrate Using Garden Hose End Sprayer	24,000	0.62	160	160

^a Worst case day (based on label) of potential exposures = mixer/loader, + application (of lawn + roses + tree + flowers), and + Post-application exposure on day of application.

Calculations:

$$\text{Residential MOE} = \frac{\text{Short-term NOEL}}{\text{Total residential exposure}}$$

$$\text{Load/Apply}_{\text{handwand}} = \frac{100 \text{ mg/kg/day}}{1 \text{ mg/kg/day}} = 100$$

$$\text{Load/Apply}_{\text{handwand}} = \frac{100 \text{ mg/kg/day}}{1.9 \text{ mg/kg/day}} = 53$$

$$\text{Load/Apply}_{\text{backpack}} = \frac{100 \text{ mg/kg/day}}{0.49 \text{ mg/kg/day}} = 200$$

$$\text{Load/Apply}_{\text{sprayer}} = \frac{100 \text{ mg/kg/day}}{0.62 \text{ mg/kg/day}} = 160$$

$$Total\ MOE = \frac{1}{\frac{1}{MOE_{food}} + \frac{1}{MOE_{residential}}}$$

$$Total\ MOE_{Using\ Handwand} = \frac{1}{\frac{1}{24,000} + \frac{1}{100}} = 100$$

$$Total\ MOE_{Using\ Handwand} = \frac{1}{\frac{1}{24,000} + \frac{1}{53}} = 53$$

$$Total\ MOE_{Using\ Backpack} = \frac{1}{\frac{1}{24,000} + \frac{1}{200}} = 200$$

$$Total\ MOE_{Using\ Sprayer} = \frac{1}{\frac{1}{24,000} + \frac{1}{160}} = 160$$

There is a potential for short-term exposure from drinking water. However, as estimated average concentrations of myclobutanil in surface and ground water are less than OPP's levels of concern for drinking water as a contribution to chronic aggregate and acute aggregate exposures, contribution to short-term exposure should not exceed OPP's levels of concern either.

RABl concludes that short-term aggregate MOEs for adults are acceptable considering the default assumptions used in the derivation of exposure estimates and the fact that a LOEL was not identified in the 28-day rat dermal toxicity study [the HDT was the NOEL in this study] used to determine the MOE. **Chemical-specific dissipation data and residential use/usage information are required to further refine these post-application exposure estimates.**

4. *Intermediate-Term Aggregate Risk*

Intermediate-term exposure scenarios are present for adults during post-application activities.

Subgroup	ARC (from DRES) (mg/kg/day)	Calculated Dietary MOE (from DRES)
U.S. Population (48 states)	0.004255	24000

Calculations:

$$\begin{aligned} \text{Dietary MOE} &= \frac{\text{Intermediate-term NOEL}}{\text{Chronic dietary exposure}} \\ &= \frac{10 \text{ mg/kg/day}}{0.004274 \text{ mg/kg/day}} = 2400 \end{aligned}$$

Subgroup	ARC (from DRES) (mg/kg/day)	Calculated Dietary MOE (from DRES)	Post-Application MOE (from Table 3)	Total MOE
U.S. Population (48 states)	0.004274	2400	250	230

Calculations:

$$\begin{aligned} \text{Total MOE} &= \frac{1}{\frac{1}{\text{MOE}_{\text{food}}} + \frac{1}{\text{MOE}_{\text{residential}}}} \\ \text{Total intermediate-term MOE} &= \frac{1}{\frac{1}{2400} + \frac{1}{250}} = 230 \end{aligned}$$

There is a potential for intermediate-term exposure from drinking water. However, as estimated average concentrations of myclobutanil in surface and ground water are less than OPP's levels of concern for drinking water as a contribution to chronic aggregate and acute aggregate exposures, contribution to intermediate-term exposure should not exceed OPP's levels of concern either.

DETERMINATION OF CANCER RISK

A cancer risk assessment is not needed since myclobutanil is classified as Category E: not carcinogenic in two acceptable animal studies.

ENDOCRINE DISRUPTOR EFFECTS

EPA is required to develop a screening program to determine whether certain substances (including all pesticides and inerts) "may have an effect in humans that is similar to an effect produced by a naturally occurring estrogen, or such other endocrine effect...." The Agency is currently working with interested stakeholders, including other government agencies, public interest groups, industry and research scientists in developing a screening and testing program and a priority setting scheme to implement this program. Congress has allowed 3 years from the passage of FQPA (August 3, 1999) to implement this program. At that time, EPA may require further testing of this active ingredient and end use products for endocrine disrupter effects.

Based on the adverse testicular findings, and increase in the number of stillborns, and a decrease in pup weight gain during lactation, in the chronic toxicity and reproduction studies in rats, myclobutanil should be considered as a candidate for evaluation as an endocrine disruptor.

DETERMINATION OF SAFETY FOR INFANTS AND CHILDREN

In assessing the potential for additional sensitivity of infants and children to residues of myclobutanil, EPA considered data from developmental toxicity studies in the rat and rabbit and a 2-generation reproductive toxicity study in the rat. The developmental toxicity studies are designed to evaluate adverse effects on the developing organism resulting from maternal pesticide exposure during prenatal development. Reproduction studies provide information relating to pre- and post-natal effects from exposure to the pesticide, information on the reproductive capability of mating animals, and data on systemic toxicity.

EPA generally defines the level of appreciable risk as exposure that is greater than 1/100 of the no observed effect level in the animal study appropriate to the particular risk assessment. This 100-fold uncertainty factor/margin of exposure (safety) is designed to account for inter-species extrapolation and intra-species variability. Under the Food Quality Protection Act (FQPA), P.L. 104-170, which was promulgated in 1996 as an amendment to the Federal Food, Drug and Cosmetic Act (FFDCA), the Agency was directed to "ensure that there is a reasonable certainty that no harm will result to infants and children" from aggregate exposure to a pesticide chemical residue. The law further states that in the case of threshold effects, for purposes of providing this reasonable certainty of no harm, "an additional tenfold margin of safety for the pesticide chemical residue and other sources of exposure shall be applied for infants and children to take into account potential pre- and post-natal toxicity and completeness of the data with respect to exposure and toxicity to infants and children. Notwithstanding such requirement for an additional margin of safety, the Administrator may use a different margin of safety for the pesticide residue only if, on the basis of reliable data, such margin will be safe for infants and children."

1. Developmental Toxicity Studies

- a. Rats. In the developmental study (MRID# 00141672) in rats, the maternal (systemic) NOEL was 93.8 mg/kg/day, based on rough hair coat, and salivation at the LOEL of 312.6 mg/kg/day. The developmental (fetal) NOEL was 93.8 mg/kg/day based on incidences of 14th rudimentary and 7th cervical ribs at the LOEL of 312.6 mg/kg/day.
- b. Rabbits. In the developmental toxicity study (MRID# 00164971) in rabbits, the maternal (systemic) NOEL was 60 mg/kg/day, based on reduced weight gain, clinical signs of toxicity and abortions at the LOEL of 200 mg/kg/day. The developmental (fetal) NOEL was 60 mg/kg/day, based on increases in number of resorptions, decreases in litter size, and a decrease in the viability index at the LOEL of 200 mg/kg/day.

2. *Reproductive Toxicity Studies*

Rats. In the 2-generation reproductive toxicity study (MRID# 00143766, 00149581) in rats, the parental (systemic) NOEL was 2.5 mg/kg/day, based on increased liver weights and liver cell hypertrophy at the LOEL of 10 mg/kg/day. The developmental (pup) NOEL was 10 mg/kg/day, based on decreased pup body weight during lactation at the LOEL of 50 mg/kg/day. The reproductive (pup) NOEL was 10 mg/kg/day, based on the increased incidence of stillborns, and atrophy of the testes, epididymides, and prostate at the LEL of 50 mg/kg/day.

3. *Pre- and Post-Natal Sensitivity*

The pre- and post-natal toxicology data base for myclobutanil is complete with respect to current toxicological data requirements. Based on the developmental and reproductive toxicity studies discussed above, for myclobutanil there does not appear to be an extra sensitivity for pre- or post-natal effects.

Based on the above, HED concludes that reliable data support use of a 100-fold margin of exposure/uncertainty factor, rather than the standard 1000-fold margin/factor, to protect infants and children.

4. *Acute Aggregate Risk for Infants and Children*

This risk assessment is not required as the HIARC did not recommend an acute dietary risk endpoint.

5. *Chronic Aggregate Risk for Infants and Children*

Using the partially refined exposure assumptions described above, HED has concluded that the percent of the RfD that will be utilized by dietary (food only) exposure to residues of myclobutanil ranges from 25% for nursing infants (<1 year old) up to 75% for non-nursing infants (<1 year old). Despite the potential for exposure to myclobutanil in drinking water, HED does not expect the chronic aggregate exposure to exceed 100% of the RfD. HED concludes that there is a reasonable certainty that no harm will result to infants and children from chronic aggregate exposure to myclobutanil residues.

6. *Short-Term Aggregate Risk for Infants and Children*

The short-term NOEL for dermal exposure is based on a dermal exposure toxicity study. Since the NOEL is based on a dermal study, oral exposures generally cannot be used directly to calculate a short-term aggregate exposure. However, as the HIARC determined that a dermal absorption factor of 100% should be used for risk assessment, oral exposures need not be multiplied by a modifying factor (converted to dermal equivalents) so that they can be compared to the dermal endpoint.

The chronic dietary exposure and calculated dietary MOE is shown below for infants (non-nursing, < 1 year old).

Subgroup	ARC (from DRES) (mg/kg/day)	Calculated Dietary MOE (from DRES)
Non-Nursing Infants (< 1 year old)	0.018836	5300

Calculations:

$$\begin{aligned} \text{Dietary MOE} &= \frac{\text{Short-term NOEL}}{\text{Chronic dietary exposure}} \\ &= \frac{100 \text{ mg/kg/day}}{0.018836 \text{ mg/kg/day}} = 5,300 \end{aligned}$$

The dermal residential exposure is 0.85 mg/kg/day (reentry). The calculated dietary MOE for non-nursing infants (<1 year old) is 5,300.

Subgroup	ARC (from DRES) (mg/kg/day)	Calculated Dietary MOE (from DRES)	Post-Application MOE (from Table 3)	Total MOE
Non-Nursing Infants (< 1 year old)	0.018836	5300	120	120

For the short-term aggregate risk of the most highly exposed subgroup (non-nursing infants (<1 year old)), the calculated MOE is 120. There is a potential for short-term exposure from drinking water. However, as estimated average concentrations of myclobutanil in surface and ground water are less than OPP's levels of concern for drinking water as a contribution to chronic aggregate and acute aggregate exposures, contribution to short-term exposure should not exceed OPP's levels of concern either. RABI concludes that short-term aggregate MOEs for non-nursing infants (<1 year old) are acceptable.

Calculations:

$$\begin{aligned} \text{Total MOE} &= \frac{1}{\frac{1}{\text{MOE}_{\text{food}}} + \frac{1}{\text{MOE}_{\text{residential}}}} \\ \text{Total short-term MOE} &= \frac{1}{\frac{1}{5,300} + \frac{1}{110}} = 110 \end{aligned}$$

There is a potential for short-term exposure from drinking water. However, as estimated average concentrations of myclobutanil in surface and ground water are less than OPP's levels of concern for drinking water as a contribution to chronic aggregate and acute aggregate exposures, contribution to short-term exposure should not exceed OPP's levels of concern either.

RABI concludes that short-term aggregate MOEs for adults are acceptable considering the default assumptions used in the derivation of exposure estimates and the fact that a LOEL was not identified in the 28-day rat dermal toxicity study [the HDT was the NOEL in this study] used to determine the MOE. **Chemical-specific dissipation data and residential use/usage information are required to further refine these post-application exposure estimates.**

7. *Intermediate-Term Aggregate Risk for Infants and Children*

The intermediate-term NOEL for dermal exposure is based on an oral exposure toxicity study. The HIARC determined that a dermal absorption factor of 100% should be used for this risk assessment.

The chronic dietary exposure from myclobutanil is 0.018836 mg/kg/day. The calculated myclobutanil dietary MOE for non-nursing infants (<1 year old) is 530.

Subgroup	ARC (from DRES) (mg/kg/day)
Non-Nursing Infants (< 1 year old)	0.018836

Calculations:

$$\begin{aligned} \text{Dietary MOE} &= \frac{\text{Intermediate-term NOEL}}{\text{Chronic dietary exposure}} \\ &= \frac{10 \text{ mg/kg/day}}{0.018836 \text{ mg/kg/day}} = 530 \end{aligned}$$

Subgroup	ARC (from DRES) (mg/kg/day)	Calculated Dietary MOE (from DRES)	Post-Application MOE (from Table 3)	Total MOE
Non-Nursing Infants (< 1 year old)	0.018836	530	160	120

The dermal residential exposure is 0.061 mg/kg/day. The calculated intermediate-term residential MOE for non-nursing infants (<1 year old) is 160.

Calculations:

$$\begin{aligned} \text{Residential MOE} &= \frac{\text{Intermediate-term NOEL}}{\text{Residential exposure}} \\ &= \frac{10 \text{ mg/kg/day}}{0.061 \text{ mg/kg/day}} = 160 \end{aligned}$$

For the intermediate-term aggregate risk of the most highly exposed subgroup (non-nursing infants (<1 year old)), the calculated MOE is 520.

Calculations:

$$\begin{aligned} \text{Total MOE} &= \frac{1}{\frac{1}{\text{MOE}_{\text{food}}} + \frac{1}{\text{MOE}_{\text{residential}}}} \\ \text{Total intermediate-term MOE} &= \frac{1}{\frac{1}{530} + \frac{1}{160}} = 120 \end{aligned}$$

There is a potential for intermediate-term exposure from drinking water. However, as estimated average concentrations of myclobutanil in surface and ground water are less than OPP's levels of concern for drinking water as a contribution to chronic aggregate and acute aggregate exposures, contribution to intermediate-term exposure should not exceed OPP's levels of concern either.

DETERMINATION OF SAFETY TO OCCUPATIONALLY EXPOSED WORKERS

1. Acute data for this formulation were available to RAB1 in conjunction with a recent import tolerance petition for bananas (PP#2E04141). The proposed work clothing and personal protective equipment (PPE) appearing on the label for Nova 40W in water-soluble pouches include long-sleeved shirt and long pants, waterproof gloves, shoes plus socks, protective eyewear and chemical-resistant headgear for overhead exposure; These work clothing and PPE are in compliance with the Worker Protection Standard(WPS).
2. Acute data for the technical are also available to RAB1. According to the recent import tolerance petition for bananas (PP#2E04141), myclobutanil is a category III for acute oral and acute dermal; category IV for primary dermal irritation and acute inhalation; and category I for primary eye irritation. Based on these values, the restricted entry interval (REI) should be 48 hours to be in compliance with the WPS. The registrant's label has a REI of 48 hours; therefore the label is in compliance.

- Occupational exposure assumptions and estimates are summarized in Tables 1 and 2, respectively.

Worker exposure estimates are based on surrogate data from the Pesticide Handlers Exposure Database (PHED) and/or the PHED Surrogate Exposure Guide (May 1997) with the worker wearing a single layer of clothing plus gloves (**note: two flagger scenarios (when there is no mechanical flagger utilized) were addressed; 1) is assumed to wear no clothing, and 2) single layer clothing with no gloves**) for myclobutanil in water-soluble pouches.

- Using these exposure assumptions, HED has concluded that the dermal MOEs that will result from the handling and application of myclobutanil by workers utilizing aerial and ground equipment, are the following ranges : *For short-term -aerial*, for mixer/loader is 4,000(for inhalation is 38,000) to 83,000 (for inhalation is 62,000) for applicator; **flagger**, for scenario 1) is 806 and 2) is 3,800 (for inhalation is 21,000); and **for groundboom**, mixer/loader is 17,000(for inhalation is 16,000) to 12,000 (for inhalation is 23,000) applicator. *For intermediate-term -aerial*, for mixer/loader is 440 to 8,300 for applicator; **flagger**, for scenario 1) is 83 and 2) is 380; and **for groundboom**, mixer/loader is 1,700 to 1,200 applicator. These MOEs do not exceed HED's level of concern, **except for the intermediate-term flagger scenario #1 (no clothing)**, for occupationally exposed workers.

OTHER CONSIDERATIONS

Metabolism in Plants

- The nature of the residue in plants is adequately understood. The residue of concern is myclobutanil plus its alcohol metabolite (free and bound), as specified in 40 CFR 180.443(a).

Analytical Enforcement Methodology

- An adequate enforcement method (Rohm and Haas Method 34S-88-10, MRID# 408033-02) is available to enforce the established tolerances. Quantitation is by GLC using an Nitrogen/Phosphorus detector for myclobutanil and an Electron Capture detector (Ni^{63}) for residues measured as the alcohol metabolite. A copy of this method is on file in PP#4E4302.

Magnitude of the Residues

- Residues of myclobutanil and its alcohol metabolite are not expected to exceed 1.0 ppm in/on peppers (bell and non-bell) as a result of this Section 18 use. **A time-limited tolerance for the combined residues of myclobutanil and its alcohol metabolite (free and bound) should be established at this level.**

Magnitude of the Residues (Meat/Milk/Poultry and Eggs)

- Secondary residues are not expected in animal commodities as no feedstuffs are associated

with these Section 18 uses. Meat/milk/poultry/egg tolerances have been established as a result of other myclobutanil uses.

Rotational Crop Restrictions

5. Information concerning the likelihood of residues in rotational crops is not available for myclobutanil. As pepper (bell and non-bell) fields are normally rotated, **HED concludes the following restriction should be added to the label for the requested Section 18: Nova[®] 40W treated fields can be rotated at any time to crops which are included on the Nova[®] 40W label. All other crops may be planted 1 year following applications of Nova[®] 40W Agricultural Fungicide.**

International Residue Limits

6. There are no Codex, Canadian or Mexican residue limits established for myclobutanil and its metabolites on the commodities included in these Section 18 requests. Thus, harmonization is not an issue for these Section 18 actions.

SUPPLEMENTAL INFORMATION

Occupational Exposure

Table 1. Occupational Exposure Assumptions

PARAMETER	ASSUMPTION
Pesticide Handlers Exposure Database (PHED), Version 1.1, Surrogate Exposure Guide (May 1997)	Mixer/Loader (wetttable powder, water soluble bags, single layer clothing plus gloves): Dermal = <u>9.8</u> µg/lb ai handled (for inhalation = 0.11 µg/lb). <i>Low Confidence Run.</i>
	Applicator - Ground (ground boom, open cab, single layer clothing plus gloves): Dermal = <u>14.0</u> µg/lb ai applied (for inhalation = 0.74 µg/lb; <i>High Confidence Run</i>). <i>Medium Confidence Run.</i>
	Applicator - Air (liquid formulations, enclosed cockpit, single layer clothing, no gloves): Dermal = <u>5.0</u> µg/lb ai applied (for inhalation = 0.07 µg/lb; <i>Medium Confidence Run</i>). <i>Low Confidence Run.</i>
	Flagger - Air (no clothing): Dermal = <u>53.0</u> µg/lb ai handled (for inhalation = 0.35 µg/lb; <i>High Confidence Run</i>). <i>Medium Confidence Run.</i>
	Flagger - Air (single layer clothing, no gloves): Dermal = <u>11.0</u> µg/lb ai handled <i>High Confidence Run.</i>
Percent Absorption	Dermal: 100% should be used for risk assessments because 1) a dermal absorption study was not available and 2) a dermal absorption factor could not be estimated due to the lack of comparative NOELs/LOELs from oral and dermal toxicity studies in the same species.
Application Type	Ground or aerial
Minimum Finish Spray	Ground: <u>20-30</u> gal/A (estimated minimum dilution rate) Air: <u>5</u> gal/A (dilution rate for mint and peppers)

PARAMETER	ASSUMPTION
Maximum Application Rate	0.4 lb ai/A
Acres Treated/Day (default values)	Ground: 88 acres Air: 350 acres
For New Mexico (Chile & Bell Peppers) average.	Based on section 18 of 30,000 Acres (maximum) statewide.
Worker Weight	60 kg (based on Tox endpoint)
Number of Farms Treated by PCO (Professional Chemical Operator)	To treat 30,000 acres: Ground: 341 days; Air: 86 days. Several operators.

Table 2. Occupational Exposure and Risk Assessment^a

Worker	Average Daily Dermal Dose ^b (mg/kg/day)	Short-Term Dermal MOE ^c	Intermediate-Term Dermal MOE ^d
Ground Mixer/Loader	5.8X10 ⁻³ (6.4X10 ⁻⁵) ^e	17,000 (16,000) ^f	1,700 (16,000) ^f
Ground Applicator	8.2X10 ⁻³ (4.3X10 ⁻⁴) ^e	12,000 (23,000) ^f	1,200 (23,000) ^f
Aerial Mixer/Loader	2.3X10 ⁻² (2.6X10 ⁻⁴) ^e	4,000 (38,000) ^f	440 (38,000) ^f
Aerial Applicator	1.2X10 ⁻³ (1.6X10 ⁻⁴) ^e	83,000 (62,000) ^f	8,300 (62,000) ^f
Flagger	1.2X10 ⁻¹ (8.2X10 ⁻⁴) ^e (2.6X10 ⁻²) ^g	806 (21,000) ^f (3,800) ^g	83 (21,000) ^f (380) ^g

^a MOEs are expressed to two significant figures.

^b Average Daily Dermal Dose (ADD) = PHED unit exposure in mg x % absorption x application rate x acres treated/day ÷ kg body weight.

^c Short-Term Occupational Dermal Exposure MOE = NOEL/ADD (where NOEL = 100 mg/kg/day).

^d Intermediate-Term Occupational Dermal Exposure MOE = NOEL/ADD (where NOEL = 10 mg/kg/day).

^e Average Daily Inhalation Dose (ADD) = PHED inhalation unit exposure in mg x % absorption x application rate x acres treated/day ÷ kg body weight.

^f Inhalation Occupational Exposure MOE = NOEL/ADD (where NOEL = 10 mg/kg/day).

^g Flagger wearing single layer of clothing, no gloves: Average Daily Dermal Dose (ADD) = PHED unit exposure in mg x % absorption x application rate x acres treated/day ÷ kg body weight.

Dietary Exposure**Table 3. Residue Consideration Summary Table**

PARAMETER	PROPOSED USE	RESIDUE DATA
CHEMICAL	Myclobutanil	Myclobutanil
FORMULATION	Nova® 40W Fungicide in Water-Soluble Pouches (Rohm and Haas, EPA Reg. No. 707-221)	Rally 60DF (Rohm and Haas, EPA Reg. No. 707-211)
CROP	Peppers (bell and non-bell)	Tomatoes
TYPE APPLICATION	Ground or aerial	ground
# APPLICATIONS	4 applications	4 or 5 applications
TIMING	10 to 14 day intervals	21 day intervals
RATE/APPLICATION	4 ounces product/A 0.1 lbs ai/A	0.063 to 0.085 lbs ai/A
RATE/YEAR or SEASON	16 ounces product/A/crop 0.4 lbs ai/A/crop	0.24 to 0.34 lbs ai/A/crop
MAXIMUM RESIDUE	N/A	0.25 ppm (4 apps @ 0.085 lbs ai/A)
RESTRICTIONS	0 day PHI	0 day PHI
RESIDUE DATA SOURCE	N/A	MRID Nos. 420192-01, -02, 423107-01 PP#1F4030/1H5616
PERFORMING LAB	N/A	Rohm and Haas

Additional Information**Progress Toward Registration.**

This is the third Section 18 request for the use of myclobutanil on peppers. IR-4 has initiated a residue program for myclobutanil on non-bell peppers conducted in 1997. These studies are expected to be completed and submitted to EPA in June 1998.

Reregistration Status.

Myclobutanil is not a FIFRA '88 reregistration active ingredient.

Attachments: Chronic DRES Analyses (3/18/98)

cc with Attachments: S. Chun (RAB1), Julianna Cruz (RAB1)
cc (Attachment only): Brian Steinwand (CEB1)
RDI: M. Lamont (5/4/98), G. Kramer (4/24/98)
S. Chun:811-Bay:CM#2:(703)305-2449:7509C:RAB1

ATTACHMENT
(Not available electronically)

TOLERANCE ASSESSMENT SYSTEM ROUTINE CHRONIC ANALYSIS

DATE: 03/10/98

PAGE: 1

CHEMICAL INFORMATION	STUDY TYPE	EFFECTS	REFERENCE DOSES	DATA GAPS/COMMENTS	STATUS
Myclobutanil (Systane/Rally) Caswell #723K CAS No. 88671-89-0 A.I. CODE: 128857 CFR No. 180.443 185.4350	2yr feeding- rat NOEL= 2.4900 mg/kg 50.00 ppm LEL= 9.8400 mg/kg 200.00 ppm ONCO: E (RfD/PR Committee)	Testicular atrophy. No evidence of carcinog- enicity in rats or mice.	ADI UF -->100 OPP RfD= 0.025000 EPA RfD= 0.000000	No data gaps.	HED reviewed 01/27/88 EPA verified 02/25/88 WHO reviewed 1992 RfD/PR reviewed 04/28/94 EPA deferred 04/28/94 On IRIS.

POPULATION SUBGROUP	TOTAL TMRC (MG/KG BODY WEIGHT/DAY)		NEW TMRC AS PERCENT OF RFD	DIFFERENCE AS PERCENT OF RFD	EFFECT OF ANTICIPATED RESIDUES	
	CURRENT TMRC*	NEW TMRC**			ARC	%RFD
U.S. POPULATION - 48 STATES	0.004967	0.005970	23.881956	4.012884	0.004274	17.09455
U.S. POPULATION - SPRING SEASON	0.004697	0.005644	22.577192	3.788332	0.004028	16.11382
U.S. POPULATION - SUMMER SEASON	0.005336	0.006353	25.412144	4.069112	0.004334	17.33722
U.S. POPULATION - FALL SEASON	0.004951	0.005972	23.888908	4.085128	0.004384	17.53524
U.S. POPULATION - WINTER SEASON	0.004864	0.005890	23.560552	4.105672	0.004324	17.29779
NORTHEAST REGION	0.005368	0.006365	25.460936	3.987504	0.004566	18.26222
NORTH CENTRAL REGION	0.005092	0.006021	24.085708	3.717304	0.004385	17.54103
SOUTHERN REGION	0.004073	0.004934	19.737572	3.444832	0.003589	14.35714
WESTERN REGION	0.005763	0.007128	28.513364	5.461904	0.004870	19.48098
HISPANICS	0.005720	0.007395	29.579900	6.700912	0.005065	20.25892
NON-HISPANIC WHITES	0.005075	0.006080	24.321500	4.021632	0.004345	17.37972
NON-HISPANIC BLACKS	0.003866	0.004485	17.941708	2.478356	0.003371	13.48485
NON-HISPANIC OTHERS	0.004996	0.006440	25.761840	5.777132	0.004441	17.76223
NURSING INFANTS (< 1 YEAR OLD)	0.009660	0.014037	56.147932	17.507704	0.006359	25.43698
NON-NURSING INFANTS (< 1 YEAR OLD)	0.025184	0.030308	121.233972	20.496200	0.018836	75.34262
FEMALES (13+ YEARS, PREGNANT)	0.003707	0.004249	16.994700	2.167480	0.003126	12.50469
FEMALES 13+ YEARS, NURSING	0.004592	0.005419	21.675828	3.309548	0.003942	15.76718
CHILDREN (1-6 YEARS OLD)	0.013163	0.016220	64.881652	12.227884	0.011492	45.96990
CHILDREN (7-12 YEARS OLD)	0.007843	0.009123	36.491708	5.119716	0.006910	27.63822
MALES (13-19 YEARS OLD)	0.004652	0.005319	21.276796	2.669700	0.004255	17.02177
FEMALES (13-19 YEARS OLD, NOT PREG. OR NURSING)	0.003855	0.004422	17.687064	2.265468	0.003415	13.66023
MALES (20 YEARS AND OLDER)	0.003066	0.003736	14.944684	2.681472	0.002693	10.77091
FEMALES (20 YEARS AND OLDER, NOT PREG. OR NURS)	0.003046	0.003692	14.766948	2.581388	0.002528	10.11002

*Current TMRC does not include new or pending tolerances.

**New TMRC includes new, pending, and published tolerances.

TOLERANCE ASSESSMENT SUMMARY FOR Myclobutanil (Systane/Rally) DATE: 03/18/98
 USING ANTICIPATED RESIDUES
 CASWELL #723K

ANALYSIS FOR POPULATION SUB-GROUP: U.S. POPULATION - 48 STATES

EXISTING ANTICIPATED RESIDUES (PUBLISHED ONLY)		
RESULT IN AN ARC OF:	0.004065	MG/KG/DAY
THE EXISTING ARC IS EQUIVALENT TO:	16.261	% OF THE ADI.
PROPOSED NEW ANTICIPATED RESIDUES (CURRENT PETITION ONLY)		
RESULT IN AN ARC OF:	0.000019	MG/KG/DAY
THESE NEW ANTICIPATED RESIDUES WILL OCCUPY:	0.075	% OF THE ADI.
IF THE NEW ANTICIPATED RESIDUES (CURRENT PETITION ONLY)		
ARE APPROVED THE RESULTANT ARC WILL BE:	0.004084	MG/KG/DAY
THE NEW ARC WILL OCCUPY	16.335	% OF THE ADI.
OTHER PENDING ANTICIPATED RESIDUES EXCLUDING THE		
CURRENT NEW PETITION HAVE AN ARC OF:	0.000190	MG/KG/DAY
THIS ARC WILL OCCUPY	0.759	% OF THE ADI.
IF ALL PENDING ANTICIPATED RESIDUES (INCLUDING THE		
CURRENT NEW PETITION) ARE GRANTED		
THE RESULTANT ARC WILL BE:	0.004274	MG/KG/DAY
THE TOTAL ARC WILL OCCUPY	17.095	% OF THE ADI.

ANALYSIS FOR POPULATION SUB-GROUP: NURSING INFANTS (< 1 YEAR OLD)

EXISTING ANTICIPATED RESIDUES (PUBLISHED ONLY)		
RESULT IN AN ARC OF:	0.005567	MG/KG/DAY
THE EXISTING ARC IS EQUIVALENT TO:	22.267	% OF THE ADI.
NO NEW ANTICIPATED RESIDUES ARE IN THE FILE.		
OTHER PENDING ANTICIPATED RESIDUES EXCLUDING THE		
CURRENT NEW PETITION HAVE AN ARC OF:	0.000793	MG/KG/DAY
THIS ARC WILL OCCUPY	3.170	% OF THE ADI.
IF ALL PENDING ANTICIPATED RESIDUES (INCLUDING THE		
CURRENT NEW PETITION) ARE GRANTED		
THE RESULTANT ARC WILL BE:	0.006359	MG/KG/DAY
THE TOTAL ARC WILL OCCUPY	25.437	% OF THE ADI.

ANALYSIS FOR POPULATION SUB-GROUP: NON-NURSING INFANTS (< 1 YEAR OLD)

EXISTING ANTICIPATED RESIDUES (PUBLISHED ONLY)		
RESULT IN AN ARC OF:	0.017906	MG/KG/DAY
THE EXISTING ARC IS EQUIVALENT TO:	71.622	% OF THE ADI.
NO NEW ANTICIPATED RESIDUES ARE IN THE FILE.		
OTHER PENDING ANTICIPATED RESIDUES EXCLUDING THE		
CURRENT NEW PETITION HAVE AN ARC OF:	0.000930	MG/KG/DAY
THIS ARC WILL OCCUPY	3.721	% OF THE ADI.
IF ALL PENDING ANTICIPATED RESIDUES (INCLUDING THE		
CURRENT NEW PETITION) ARE GRANTED		
THE RESULTANT ARC WILL BE:	0.018836	MG/KG/DAY
THE TOTAL ARC WILL OCCUPY	75.343	% OF THE ADI.

TOLERANCE ASSESSMENT SUMMARY FOR Myclobutanil (Systane/Rally) DATE: 03/10/98
 USING ANTICIPATED RESIDUES
 CASWELL #723K

ANALYSIS FOR POPULATION SUB-GROUP: CHILDREN (1-6 YEARS OLD)

EXISTING ANTICIPATED RESIDUES (PUBLISHED ONLY)		
RESULT IN AN ARC OF:	0.010897	MG/KG/DAY
THE EXISTING ARC IS EQUIVALENT TO:	43.587	% OF THE ADI.
PROPOSED NEW ANTICIPATED RESIDUES (CURRENT PETITION ONLY)		
RESULT IN AN ARC OF:	<0.000001	MG/KG/DAY
THESE NEW ANTICIPATED RESIDUES WILL OCCUPY:	0.000	% OF THE ADI.
IF THE NEW ANTICIPATED RESIDUES (CURRENT PETITION ONLY)		
ARE APPROVED THE RESULTANT ARC WILL BE:	0.010897	MG/KG/DAY
THE NEW ARC WILL OCCUPY	43.587	% OF THE ADI.
OTHER PENDING ANTICIPATED RESIDUES EXCLUDING THE		
CURRENT NEW PETITION HAVE AN ARC OF:	0.000596	MG/KG/DAY
THIS ARC WILL OCCUPY	2.383	% OF THE ADI.
IF ALL PENDING ANTICIPATED RESIDUES (INCLUDING THE		
CURRENT NEW PETITION) ARE GRANTED		
THE RESULTANT ARC WILL BE:	0.011492	MG/KG/DAY
THE TOTAL ARC WILL OCCUPY	45.970	% OF THE ADI.

ANALYSIS FOR POPULATION SUB-GROUP: CHILDREN (7-12 YEARS OLD)

EXISTING ANTICIPATED RESIDUES (PUBLISHED ONLY)		
RESULT IN AN ARC OF:	0.006664	MG/KG/DAY
THE EXISTING ARC IS EQUIVALENT TO:	26.655	% OF THE ADI.
NO NEW ANTICIPATED RESIDUES ARE IN THE FILE.		
OTHER PENDING ANTICIPATED RESIDUES EXCLUDING THE		
CURRENT NEW PETITION HAVE AN ARC OF:	0.000246	MG/KG/DAY
THIS ARC WILL OCCUPY	0.983	% OF THE ADI.
IF ALL PENDING ANTICIPATED RESIDUES (INCLUDING THE		
CURRENT NEW PETITION) ARE GRANTED		
THE RESULTANT ARC WILL BE:	0.006910	MG/KG/DAY
THE TOTAL ARC WILL OCCUPY	27.638	% OF THE ADI.

ANTICIPATED RESIDUE INFORMATION FOR CASWELL NUMBER 723K

DATE: 03/18/98

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CHEMICAL	STUDY TYPE	EFFECTS	REFERENCE DOSES	DATA GAPS/COMMENTS	STATUS
Myclobutanil (Systane/Rally) Caswell #723K CAS No. 88671-89-0 A.I. CODE: 128857 CFR No. 180.443 185.4350	2yr feeding- rat NOEL= 2.4900 mg/kg 50.00 ppm LEL= 9.8400 mg/kg 200.00 ppm ONCO: E (RfD/PR Committee)	Testicular atrophy. No evidence of carcinog- enicity in rats or mice.	ADI UF -->100 OPP RfD= 0.025000 EPA RfD= 0.000000	No data gaps.	NED reviewed 01/27/88 EPA verified 02/25/88 WHO reviewed 1992 RfD/PR reviewed 04/28/94 EPA deferred 04/28/94 On IRIS.

FOOD CODE	FOOD	FOOD FORM	PET.#	TOLERANCE (ppm)	ANTICIPATED RESIDUE (ppm)	AR STATISTIC TYPE	% CROP TREATED	RES. VALUE USED IN TAS RUN (ppm)
01014AA	GRAPES-FRESH	10 RAW-FRESH OR NFS	7F3476	P 1.000000	1.000000		79.00	0.790000
01014AA	GRAPES-FRESH	21 COOKED-NFS	7F3476	P 1.000000	1.000000		79.00	0.790000
01014AA	GRAPES-FRESH	31 COOKED-FRESH OR CANNED	7F3476	P 1.000000	1.000000		79.00	0.790000
01014DA	GRAPES-RAISINS	10 RAW-FRESH OR NFS	7H5524	P 10.000000	10.000000C		79.00	7.900000
01014DA	GRAPES-RAISINS	21 COOKED-NFS	7H5524	P 10.000000	10.000000C		79.00	7.900000
01014DA	GRAPES-RAISINS	22 COOKED-FRESH-BAKED	7H5524	P 10.000000	10.000000C		79.00	7.900000
01014JA	GRAPES-JUICE	10 RAW-FRESH OR NFS	7F3476	P 1.000000	1.000000		79.00	0.790000
01014JA	GRAPES-JUICE	15 RAW-FRESH OR CANNED	7F3476	P 1.000000	1.000000		79.00	0.790000
01014JA	GRAPES-JUICE	21 COOKED-NFS	7F3476	P 1.000000	1.000000		79.00	0.790000
01016AA	STRAWBERRIES	10 RAW-FRESH OR NFS	97FL001	P 0.500000	0.500000		100.00	0.500000
01016AA	STRAWBERRIES	21 COOKED-NFS	97FL001	P 0.500000	0.500000		100.00	0.500000
01016AA	STRAWBERRIES	70 RAW-FROZEN	97FL001	P 0.500000	0.500000		100.00	0.500000
03001AA	ALMONDS	10 RAW-FRESH OR NFS	0F3876	P 0.100000	0.100000		1.00	0.001000
03001AA	ALMONDS	21 COOKED-NFS	0F3876	P 0.100000	0.100000		1.00	0.001000
03001AA	ALMONDS	22 COOKED-FRESH-BAKED	0F3876	P 0.100000	0.100000		1.00	0.001000
04001AA	APPLES-FRESH	10 RAW-FRESH OR NFS	7F3476	P 0.500000	0.500000		60.00	0.300000
04001AA	APPLES-FRESH	21 COOKED-NFS	7F3476	P 0.500000	0.500000		60.00	0.300000
04001AA	APPLES-FRESH	31 COOKED-FRESH OR CANNED	7F3476	P 0.500000	0.500000		60.00	0.300000
04001AA	APPLES-FRESH	62 COOKED-FRESH OR FROZEN-BAKED	7F3476	P 0.500000	0.500000		60.00	0.300000
04001DA	APPLES-DRIED	10 RAW-FRESH OR NFS	7F3476	P 0.500000	0.500000		60.00	0.300000
04001DA	APPLES-DRIED	22 COOKED-FRESH-BAKED	7F3476	P 0.500000	0.500000		60.00	0.300000
04001DA	APPLES-DRIED	62 COOKED-FRESH OR FROZEN-BAKED	7F3476	P 0.500000	0.500000		60.00	0.300000
04001JA	APPLES-JUICE	15 RAW-FRESH OR CANNED	7F3476	P 0.500000	0.500000		60.00	0.300000
04001JA	APPLES-JUICE	31 COOKED-FRESH OR CANNED	7F3476	P 0.500000	0.500000		60.00	0.300000
04002AA	CRABAPPLES	00 NOT SPECIFIED (NO CONSUMPTION)	9F3812	A 0.500000	0.500000		100.00	0.500000
04003AA	PEARS-FRESH	10 RAW-FRESH OR NFS	9F3812	A 0.500000	0.500000		8.00	0.040000
04003AA	PEARS-FRESH	31 COOKED-FRESH OR CANNED	9F3812	A 0.500000	0.500000		8.00	0.040000
04003AA	PEARS-FRESH	51 COOKED-CANNED	9F3812	A 0.500000	0.500000		8.00	0.040000
04003AA	PEARS-FRESH	62 COOKED-FRESH OR FROZEN-BAKED	9F3812	A 0.500000	0.500000		8.00	0.040000
04003DA	PEARS-DRIED	10 RAW-FRESH OR NFS	9F3812	A 0.500000	0.500000		8.00	0.040000
04003DA	PEARS-DRIED	21 COOKED-NFS	9F3812	A 0.500000	0.500000		8.00	0.040000
04004AA	QUINCES	00 NOT SPECIFIED (NO CONSUMPTION)	9F3812	A 0.500000	0.500000		100.00	0.500000
05001AA	APRICOTS-FRESH	10 RAW-FRESH OR NFS	1F3954	P 2.000000	2.000000		1.00	0.020000
05001AA	APRICOTS-FRESH	21 COOKED-NFS	1F3954	P 2.000000	2.000000		1.00	0.020000
05001AA	APRICOTS-FRESH	31 COOKED-FRESH OR CANNED	1F3954	P 2.000000	2.000000		1.00	0.020000
05001DA	APRICOTS-DRIED	10 RAW-FRESH OR NFS	1F3954	P 2.000000	2.000000		1.00	0.020000
05001DA	APRICOTS-DRIED	22 COOKED-FRESH-BAKED	1F3954	P 2.000000	2.000000		1.00	0.020000
05002AA	CHERRIES-FRESH	10 RAW-FRESH OR NFS	2F4116	P 5.000000	5.000000		47.00	2.350000
05002AA	CHERRIES-FRESH	21 COOKED-NFS	2F4116	P 5.000000	5.000000		47.00	2.350000
05002AA	CHERRIES-FRESH	31 COOKED-FRESH OR CANNED	2F4116	P 5.000000	5.000000		47.00	2.350000

ANTICIPATED RESIDUE INFORMATION FOR CASWELL NUMBER 723K

DATE: 03/19/98

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CHEMICAL	STUDY TYPE	EFFECTS	REFERENCE DOSES	DATA GAPS/COMMENTS	STATUS
Myclobutanil (Systane/Rally) Caswell #723K CAS No. 88671-89-0 A.I. CODE: 128857 CFR No. 180.443 185.4350	2yr feeding- rat NOEL= 2.4900 mg/kg 50.00 ppm LEL= 9.8400 mg/kg 200.00 ppm ONCO: E (RfD/PR Committee)	Testicular atrophy. No evidence of carcinogenicity in rats or mice.	ADI UF -->100 OPP RfD= 0.025000 EPA RfD= 0.000000	No data gaps.	HED reviewed 01/27/88 EPA verified 02/25/88 WHO reviewed 1992 RfD/PR reviewed 04/28/94 EPA deferred 04/28/94 On IRIS.

FOOD CODE	FOOD	FOOD FORM	PET.#	TOLERANCE (ppm)	ANTICIPATED RESIDUE (ppm)	AR STATISTIC TYPE	% CROP TREATED	RES. VALUE USED IN TAS RUN (ppm)
05002AA	CHERRIES-FRESH	62 COOKED-FRESH OR FROZEN-BAKED	2F4116	P 5.000000	5.000000		47.00	2.350000
05002DA	CHERRIES-DRIED	00 NOT SPECIFIED (NO CONSUMPTION)	2F4116	P 5.000000	5.000000		47.00	2.350000
05002JA	CHERRIES-JUICE	15 RAW-FRESH OR CANNED	2F4116	P 5.000000	5.000000		47.00	2.350000
05002JA	CHERRIES-JUICE	21 COOKED-NFS	2F4116	P 5.000000	5.000000		47.00	2.350000
05003AA	NECTARINES	10 RAW-FRESH OR NFS	9F3811	P 2.000000	2.000000		21.00	0.420000
05004AA	PEACHES-FRESH	10 RAW-FRESH OR NFS	9F3811	P 2.000000	2.000000		22.00	0.440000
05004AA	PEACHES-FRESH	21 COOKED-NFS	9F3811	P 2.000000	2.000000		22.00	0.440000
05004AA	PEACHES-FRESH	31 COOKED-FRESH OR CANNED	9F3811	P 2.000000	2.000000		22.00	0.440000
05004AA	PEACHES-FRESH	51 COOKED-CANNED	9F3811	P 2.000000	2.000000		22.00	0.440000
05004DA	PEACHES-DRIED	10 RAW-FRESH OR NFS	9F3811	P 2.000000	2.000000		22.00	0.440000
05004DA	PEACHES-DRIED	21 COOKED-NFS	9F3811	P 2.000000	2.000000		22.00	0.440000
05005AA	PLUMS-FRESH	10 RAW-FRESH OR NFS	1F3954	P 2.000000	2.000000		3.00	0.060000
05005AA	PLUMS-FRESH	31 COOKED-FRESH OR CANNED	1F3954	P 2.000000	2.000000		3.00	0.060000
05005DA	PLUMS-PRUNES	10 RAW-FRESH OR NFS	1H5608	P 8.000000	8.000000C		3.00	0.240000
05005DA	PLUMS-PRUNES	21 COOKED-NFS	1H5608	P 8.000000	8.000000C		3.00	0.240000
05005DA	PLUMS-PRUNES	31 COOKED-FRESH OR CANNED	1H5608	P 8.000000	8.000000C		3.00	0.240000
05005JA	PRUNE-JUICE	10 RAW-FRESH OR NFS	1F3954	P 2.000000	2.000000		3.00	0.060000
05005JA	PRUNE-JUICE	62 COOKED-FRESH OR FROZEN-BAKED	1F3954	P 2.000000	2.000000		3.00	0.060000
06002AA	BANANAS-UNSPEC	22 COOKED-FRESH-BAKED	2E04141	A 4.000000	0.800000		100.00	0.800000
06002AB	BANANAS-FRESH	10 RAW-FRESH OR NFS	2E04141	A 4.000000	0.800000		100.00	0.800000
06002AB	BANANAS-FRESH	21 COOKED-NFS	2E04141	A 4.000000	0.800000		100.00	0.800000
06002AB	BANANAS-FRESH	31 COOKED-FRESH OR CANNED	2E04141	A 4.000000	0.800000		100.00	0.800000
06002DA	BANANAS-DRIED	10 RAW-FRESH OR NFS	2E04141	A 4.000000	0.800000		100.00	0.800000
06002DA	BANANAS-DRIED	21 COOKED-NFS	2E04141	A 4.000000	0.800000		100.00	0.800000
06016AA	PLANTAINS	21 COOKED-NFS	2E04141	A 4.000000	0.800000		100.00	0.800000
06016AA	PLANTAINS	23 COOKED-FRESH-BOILED	2E04141	A 4.000000	0.800000		100.00	0.800000
06016AA	PLANTAINS	25 COOKED-FRESH-FRIED	2E04141	A 4.000000	0.800000		100.00	0.800000
08020AA	HOPS	21 COOKED-NFS	98WA006	N 5.000000	5.000000		100.00	5.000000
10002AA	CANTALOUPE-UNSP	00 NOT SPECIFIED (NO CONSUMPTION)	SECT18	P 0.300000	0.300000		100.00	0.300000
10002AB	CANTALOUPE-PULP	10 RAW-FRESH OR NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10002AB	CANTALOUPE-PULP	21 COOKED-NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10003AA	CASABAS	10 RAW-FRESH OR NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10004AA	CRENSHAW	00 NOT SPECIFIED (NO CONSUMPTION)	SECT18	P 0.300000	0.300000		100.00	0.300000
10005AA	HONEYDEW MELONS	10 RAW-FRESH OR NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10007AA	PERSION MELONS	00 NOT SPECIFIED (NO CONSUMPTION)	SECT18	P 0.300000	0.300000		100.00	0.300000
10008AA	WATERMELON	10 RAW-FRESH OR NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10008AA	WATERMELON	21 COOKED-NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10010AA	CUCUMBERS	10 RAW-FRESH OR NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10010AA	CUCUMBERS	11 RAW-FRESH-PICKLED,CORNED,OR CURED	SECT18	P 0.300000	0.300000		100.00	0.300000
10010AA	CUCUMBERS	21 COOKED-NFS	SECT18	P 0.300000	0.300000		100.00	0.300000

ANTICIPATED RESIDUE INFORMATION FOR CASWELL NUMBER 723K

DATE: 03/18/98

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CHEMICAL	STUDY TYPE	EFFECTS	REFERENCE DOSES	DATA GAPS/COMMENTS	STATUS
Myclobutanil (Systane/Rally) Caswell #723K CAS No. 88671-89-0 A.I. CODE: 128857 CFR No. 180.443 185.4350	2yr feeding- rat NOEL= 2.4900 mg/kg 50.00 ppm LEL= 9.8400 mg/kg 200.00 ppm OMCO: E (RfD/PR Committee)	Testicular atrophy. No evidence of carcinog- enicity in rats or mice.	ADI UF -->100 OPP RfD= 0.025000 EPA RfD= 0.000000	No data gaps.	HED reviewed 01/27/88 EPA verified 02/25/88 WHO reviewed 1992 RfD/PR reviewed 04/28/94 EPA deferred 04/28/94 On IRIS.

FOOD CODE	FOOD	FOOD FORM	PET.#	TOLERANCE (ppm)	ANTICIPATED RESIDUE (ppm)	AR STATISTIC TYPE	% CROP TREATED	RES. VALUE USED IN TAS RUN (ppm)
10011AA	PUMPKIN	21 COOKED-NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10011AA	PUMPKIN	22 COOKED-FRESH-BAKED	SECT18	P 0.300000	0.300000		100.00	0.300000
10011AA	PUMPKIN	62 COOKED-FRESH OR FROZEN-BAKED	SECT18	P 0.300000	0.300000		100.00	0.300000
10013AA	SQUASH-SUMMER	10 RAW-FRESH OR NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10013AA	SQUASH-SUMMER	21 COOKED-NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10014AA	SQUASH-WINTER	10 RAW-FRESH OR NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10014AA	SQUASH-WINTER	21 COOKED-NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10014AA	SQUASH-WINTER	31 COOKED-FRESH OR CANNED	SECT18	P 0.300000	0.300000		100.00	0.300000
10017AA	BITTER MELON	21 COOKED-NFS	SECT18	P 0.300000	0.300000		100.00	0.300000
10020AA	TOMELGOURD	00 NOT SPECIFIED (NO CONSUMPTION)	SECT18	P 0.300000	0.300000		100.00	0.300000
11003AA	PEPPERS,SWEET	10 RAW-FRESH OR NFS TLT 7/1/98	97CA036	P 1.000000	1.000000		100.00	1.000000
11003AA	PEPPERS,SWEET	21 COOKED-NFS "	97CA036	P 1.000000	1.000000		100.00	1.000000
11003AB	CHILLI PEPPERS	00 NOT SPECIFIED (NO CONSUMPTION) "	97CA036	P 1.000000	1.000000		100.00	1.000000
11003AD	PEPPERS-OTHER	10 RAW-FRESH OR NFS "	97CA036	P 1.000000	1.000000		100.00	1.000000
11003AD	PEPPERS-OTHER	21 COOKED-NFS "	97CA036	P 1.000000	1.000000		100.00	1.000000
11003AD	PEPPERS-OTHER	51 COOKED-CANNED "	97CA036	P 1.000000	1.000000		100.00	1.000000
11004AA	PIMIENTOS	10 RAW-FRESH OR NFS "	97CA036	P 1.000000	1.000000		100.00	1.000000
11004AA	PIMIENTOS	21 COOKED-NFS "	97CA036	P 1.000000	1.000000		100.00	1.000000
11004AA	PIMIENTOS	31 COOKED-FRESH OR CANNED "	97CA036	P 1.000000	1.000000		100.00	1.000000
11005AA	TOMATOES-WHOLE	10 RAW-FRESH OR NFS TLT 7/28/98	97CA042	P 0.300000	0.300000		100.00	0.300000
11005AA	TOMATOES-WHOLE	21 COOKED-NFS "	97CA042	P 0.300000	0.300000		100.00	0.300000
11005AA	TOMATOES-WHOLE	31 COOKED-FRESH OR CANNED "	97CA042	P 0.300000	0.300000		100.00	0.300000
11005JA	TOMATOES-JUICE	10 RAW-FRESH OR NFS "	97CA042	P 0.300000	0.300000		100.00	0.300000
11005JA	TOMATOES-JUICE	21 COOKED-NFS "	97CA042	P 0.300000	0.300000		100.00	0.300000
11005RA	TOMATOES-PUREE	10 RAW-FRESH OR NFS "	97CA042	P 0.600000	0.600000		100.00	0.600000
11005RA	TOMATOES-PUREE	21 COOKED-NFS "	97CA042	P 0.600000	0.600000		100.00	0.600000
11005RA	TOMATOES-PUREE	31 COOKED-FRESH OR CANNED "	97CA042	P 0.600000	0.600000		100.00	0.600000
11005RA	TOMATOES-PUREE	32 COOKED-FRESH OR CANNED-BAKED "	97CA042	P 0.600000	0.600000		100.00	0.600000
11005RA	TOMATOES-PUREE	51 COOKED-CANNED "	97CA042	P 0.600000	0.600000		100.00	0.600000
11005TA	TOMATOES-PASTE	21 COOKED-NFS "	97CA042	P 1.200000	1.200000		100.00	1.200000
11005TA	TOMATOES-PASTE	22 COOKED-FRESH-BAKED "	97CA042	P 1.200000	1.200000		100.00	1.200000
11005TA	TOMATOES-PASTE	31 COOKED-FRESH OR CANNED "	97CA042	P 1.200000	1.200000		100.00	1.200000
11005UA	TOMATOES-CATSUP	21 COOKED-NFS "	97CA042	P 0.600000	0.600000		100.00	0.600000
16002AA	ASPARAGUS	21 COOKED-NFS	97CA026	P 0.010000	0.010000		100.00	0.010000
16002AA	ASPARAGUS	23 COOKED-FRESH-BOILED	97CA026	P 0.010000	0.010000		100.00	0.010000
270030A	COTTONSEED-OIL	18 PROCESSED OIL	4F4317	P 0.020000	0.020000		1.00	0.000200
27003WA	COTTONSEED-MEAL	18 PROCESSED OIL	4F4317	P 0.020000	0.020000		1.00	0.000200
28080AA	PEPPERMINT	00 NOT SPECIFIED (NO CONSUMPTION) 7/1/98	97ID014	P 2.500000	2.500000		100.00	2.500000
28080QA	PEPPERMINT-OIL	00 NOT SPECIFIED (NO CONSUMPTION) "	97ID014	P 2.500000	2.500000		100.00	2.500000
28081AA	SPEARMINT	00 NOT SPECIFIED (NO CONSUMPTION) "	97ID014	P 2.500000	2.500000		100.00	2.500000

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CHEMICAL	STUDY TYPE	EFFECTS	REFERENCE DOSES	DATA GAPS/COMMENTS	STATUS
Myclobutanil (Systane/Rally) Caswell #723K CAS No. 88671-89-0 A.I. CODE: 128857 CFR No. 180.443 185.4350	2yr feeding- rat NOEL= 2.4900 mg/kg 50.00 ppm LEL= 9.8400 mg/kg 200.00 ppm ONCO: E (RfD/PR Committee)	Testicular atrophy. No evidence of carcinog- enicity in rats or mice.	ADI UF -->100 OPP RfD= 0.025000 EPA RfD= 0.000000	No data gaps.	HED reviewed 01/27/88 EPA verified 02/25/88 WHO reviewed 1992 RfD/PR reviewed 04/28/94 EPA deferred 04/28/94 On IRIS.

FOOD CODE	FOOD	FOOD FORM	PET.#	TOLERANCE (ppm)	ANTICIPATED RESIDUE (ppm)	AR STATISTIC TYPE	% CROP TREATED	RES. VALUE USED IN TAS RUN (ppm)
280810A	SPEARMINT-OIL	00 NOT SPECIFIED (NO CONSUMPTION) "	971D014	P 2.500000	2.500000		100.00	2.500000
43058AA	WINE AND SHERRY	10 RAW-FRESH OR NFS	7F3476	P 1.000000	1.000000		79.00	0.790000
43058AA	WINE AND SHERRY	21 COOKED-NFS	7F3476	P 1.000000	1.000000		79.00	0.790000
50000DB	MILK-NON-FAT SOL	10 RAW-FRESH OR NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
50000DB	MILK-NON-FAT SOL	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
50000DB	MILK-NON-FAT SOL	51 COOKED-CANNED	0F3876	P 0.200000	0.200000		100.00	0.200000
50000FA	MILK-FAT SOLIDS	10 RAW-FRESH OR NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
50000FA	MILK-FAT SOLIDS	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
50000FA	MILK-FAT SOLIDS	51 COOKED-CANNED	0F3876	P 0.200000	0.200000		100.00	0.200000
50000SA	MILK SUG (LACT)	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
50000SA	MILK SUG (LACT)	51 COOKED-CANNED	0F3876	P 0.200000	0.200000		100.00	0.200000
53001BA	BEEF-MEAT BYP	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
53001BA	BEEF-MEAT BYP	26 COOKED-FRESH-PICKLED, CORNED, OR CURED	0F3876	P 0.200000	0.200000		100.00	0.200000
53001BB	BEEF-OTH ORGAN	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
53001BB	BEEF-OTH ORGAN	51 COOKED-CANNED	0F3876	P 0.200000	0.200000		100.00	0.200000
53001DA	BEEF-DRIED	21 COOKED-NFS	0F3876	P 0.100000	0.100000		100.00	0.100000
53001FA	BEEF-FAT	10 RAW-FRESH OR NFS	0F3876	P 0.050000	0.050000		100.00	0.050000
53001FA	BEEF-FAT	21 COOKED-NFS	0F3876	P 0.050000	0.050000		100.00	0.050000
53001FA	BEEF-FAT	22 COOKED-FRESH-BAKED	0F3876	P 0.050000	0.050000		100.00	0.050000
53001FA	BEEF-FAT	23 COOKED-FRESH-BOILED	0F3876	P 0.050000	0.050000		100.00	0.050000
53001FA	BEEF-FAT	24 COOKED-FRESH-BROILED	0F3876	P 0.050000	0.050000		100.00	0.050000
53001FA	BEEF-FAT	25 COOKED-FRESH-FRIED	0F3876	P 0.050000	0.050000		100.00	0.050000
53001KA	BEEF-KIDNEY	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
53001LA	BEEF-LIVER	25 COOKED-FRESH-FRIED	0F3876	P 1.000000	1.000000		100.00	1.000000
53001LA	BEEF-LIVER	31 COOKED-FRESH OR CANNED	0F3876	P 1.000000	1.000000		100.00	1.000000
53001MA	BEEF-LEAN	10 RAW-FRESH OR NFS	0F3876	P 0.100000	0.100000		100.00	0.100000
53001MA	BEEF-LEAN	21 COOKED-NFS	0F3876	P 0.100000	0.100000		100.00	0.100000
53001MA	BEEF-LEAN	22 COOKED-FRESH-BAKED	0F3876	P 0.100000	0.100000		100.00	0.100000
53001MA	BEEF-LEAN	23 COOKED-FRESH-BOILED	0F3876	P 0.100000	0.100000		100.00	0.100000
53001MA	BEEF-LEAN	24 COOKED-FRESH-BROILED	0F3876	P 0.100000	0.100000		100.00	0.100000
53002BA	GOAT-MEAT BYP	00 NOT SPECIFIED (NO CONSUMPTION)	0F3876	P 0.200000	0.200000		100.00	0.200000
53002BB	GOAT-OTH ORGAN	00 NOT SPECIFIED (NO CONSUMPTION)	0F3876	P 0.200000	0.200000		100.00	0.200000
53002FA	GOAT-FAT	23 COOKED-FRESH-BOILED	0F3876	P 0.050000	0.050000		100.00	0.050000
53002FA	GOAT-FAT	25 COOKED-FRESH-FRIED	0F3876	P 0.050000	0.050000		100.00	0.050000
53002KA	GOAT-KIDNEY	00 NOT SPECIFIED (NO CONSUMPTION)	0F3876	P 0.200000	0.200000		100.00	0.200000
53002LA	GOAT-LIVER	00 NOT SPECIFIED (NO CONSUMPTION)	0F3876	P 1.000000	1.000000		100.00	1.000000
53002MA	GOAT-LEAN	23 COOKED-FRESH-BOILED	0F3876	P 0.100000	0.100000		100.00	0.100000
53002MA	GOAT-LEAN	25 COOKED-FRESH-FRIED	0F3876	P 0.100000	0.100000		100.00	0.100000
53003AA	HORSE	00 NOT SPECIFIED (NO CONSUMPTION)	0F3876	P 1.000000	1.000000		100.00	1.000000
53005BA	SHEEP-MEAT BYP	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000

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CHEMICAL	STUDY TYPE	EFFECTS	REFERENCE DOSES	DATA GAPS/COMMENTS	STATUS
Myclobutanil (Systane/Rally) Caswell #723K CAS No. 88671-89-0 A.I. CODE: 128857 CFR No. 180.443 185.4350	2yr feeding- rat MOEL= 2.4900 mg/kg 50.00 ppm LEL= 9.8400 mg/kg 200.00 ppm ONCO: E (RfD/PR Committee)	Testicular atrophy. No evidence of carcinog- enicity in rats or mice.	ADI UF -->100 OPP RfD= 0.025000 EPA RfD= 0.000000	No data gaps.	HED reviewed 01/27/88 EPA verified 02/25/88 WHO reviewed 1992 RfD/PR reviewed 04/28/94 EPA deferred 04/28/94 On IRIS.

FOOD CODE	FOOD	FOOD FORM	PET.#	TOLERANCE (ppm)	ANTICIPATED RESIDUE (ppm)	AR STATISTIC TYPE	% CROP TREATED	RES. VALUE USED IN TAS RUN (ppm)
53005BB	SHEEP-OTH ORGAN	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
53005FA	SHEEP-FAT	21 COOKED-NFS	0F3876	P 0.050000	0.050000		100.00	0.050000
53005KA	SHEEP-KIDNEY	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
53005LA	SHEEP-LIVER	00 NOT SPECIFIED (NO CONSUMPTION)	0F3876	P 1.000000	1.000000		100.00	1.000000
53005MA	SHEEP-LEAN	21 COOKED-NFS	0F3876	P 0.100000	0.100000		100.00	0.100000
53005MA	SHEEP-LEAN	31 COOKED-FRESH OR CANNED	0F3876	P 0.100000	0.100000		100.00	0.100000
53006BA	PORK-MEAT BYP	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
53006BB	PORK-OTH ORGAN	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
53006BB	PORK-OTH ORGAN	26 COOKED-FRESH-PICKLED,CORNE D,OR CURED	0F3876	P 0.200000	0.200000		100.00	0.200000
53006FA	PORK-FAT	10 RAW-FRESH OR NFS	0F3876	P 0.050000	0.050000		100.00	0.050000
53006FA	PORK-FAT	21 COOKED-NFS	0F3876	P 0.050000	0.050000		100.00	0.050000
53006FA	PORK-FAT	23 COOKED-FRESH-BOILED	0F3876	P 0.050000	0.050000		100.00	0.050000
53006FA	PORK-FAT	25 COOKED-FRESH-FRIED	0F3876	P 0.050000	0.050000		100.00	0.050000
53006FA	PORK-FAT	26 COOKED-FRESH-PICKLED,CORNE D,OR CURED	0F3876	P 0.050000	0.050000		100.00	0.050000
53006KA	PORK-KIDNEY	21 COOKED-NFS	0F3876	P 0.200000	0.200000		100.00	0.200000
53006LA	PORK-LIVER	21 COOKED-NFS	0F3876	P 1.000000	1.000000		100.00	1.000000
53006LA	PORK-LIVER	25 COOKED-FRESH-FRIED	0F3876	P 1.000000	1.000000		100.00	1.000000
53006MA	PORK-LEAN	21 COOKED-NFS	0F3876	P 0.100000	0.100000		100.00	0.100000
53006MA	PORK-LEAN	25 COOKED-FRESH-FRIED	0F3876	P 0.100000	0.100000		100.00	0.100000
53006MA	PORK-LEAN	26 COOKED-FRESH-PICKLED,CORNE D,OR CURED	0F3876	P 0.100000	0.100000		100.00	0.100000
55008BA	TURKEY-BYP	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55008BA	TURKEY-BYP	26 COOKED-FRESH-PICKLED,CORNE D,OR CURED	7F3476	P 0.020000	0.020000		100.00	0.020000
55008LA	TURKEY ORGAN	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55008LA	TURKEY ORGAN	25 COOKED-FRESH-FRIED	7F3476	P 0.020000	0.020000		100.00	0.020000
55008MA	TURKEY W/O SKIN	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55008MA	TURKEY W/O SKIN	31 COOKED-FRESH OR CANNED	7F3476	P 0.020000	0.020000		100.00	0.020000
55008MA	TURKEY W/O SKIN	62 COOKED-FRESH OR FROZEN-BAKED	7F3476	P 0.020000	0.020000		100.00	0.020000
55008MB	TURKEY+SKIN	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55008MB	TURKEY+SKIN	25 COOKED-FRESH-FRIED	7F3476	P 0.020000	0.020000		100.00	0.020000
55008MC	TURKEY-UNSPEC	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55013BA	POULTRY,OTH-BYP	00 NOT SPECIFIED (NO CONSUMPTION)	7F3476	P 0.020000	0.020000		100.00	0.020000
55013LA	POULTRY,ORGAN	25 COOKED-FRESH-FRIED	7F3476	P 0.020000	0.020000		100.00	0.020000
55013MA	POULTRY,OTHER	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AA	EGGS-WHOLE	10 RAW-FRESH OR NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AA	EGGS-WHOLE	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AA	EGGS-WHOLE	22 COOKED-FRESH-BAKED	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AA	EGGS-WHOLE	23 COOKED-FRESH-BOILED	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AA	EGGS-WHOLE	25 COOKED-FRESH-FRIED	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AB	EGGS-WHITE ONLY	10 RAW-FRESH OR NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AB	EGGS-WHITE ONLY	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000

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CHEMICAL	STUDY TYPE	EFFECTS	REFERENCE DOSES	DATA GAPS/COMMENTS	STATUS
Myclobutanil (Systane/Rally) Caswell #723K CAS No. 88671-89-0 A.I. CODE: 128857 CFR No. 180.443 185.4350	2yr feeding- rat NOEL= 2.4900 mg/kg 50.00 ppm LEL= 9.8400 mg/kg 200.00 ppm ONCO: E (Rfd/PR Committee)	Testicular atrophy. No evidence of carcinog- enicity in rats or mice.	ADI UF -->100 OPP Rfd= 0.025000 EPA Rfd= 0.000000	No data gaps.	HED reviewed 01/27/88 EPA verified 02/25/88 WHO reviewed 1992 Rfd/PR reviewed 04/28/94 EPA deferred 04/28/94 On IRIS.

FOOD CODE	FOOD	FOOD FORM	PET.#	TOLERANCE (ppm)	ANTICIPATED RESIDUE (ppm)	AR STATISTIC TYPE	% CROP TREATED	RES. VALUE USED IN TAS RUN (ppm)
55014AB	EGGS-WHITE ONLY	22 COOKED-FRESH-BAKED	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AB	EGGS-WHITE ONLY	62 COOKED-FRESH OR FROZEN-BAKED	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AB	EGGS-WHITE ONLY	81 COOKED-FROZEN	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AC	EGGS-YOLK ONLY	10 RAW-FRESH OR NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AC	EGGS-YOLK ONLY	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AC	EGGS-YOLK ONLY	25 COOKED-FRESH-FRIED	7F3476	P 0.020000	0.020000		100.00	0.020000
55014AC	EGGS-YOLK ONLY	31 COOKED-FRESH OR CANNED	7F3476	P 0.020000	0.020000		100.00	0.020000
55015BA	CHICKEN-BYP	00 NOT SPECIFIED (NO CONSUMPTION)	7F3476	P 0.020000	0.020000		100.00	0.020000
55015LA	CHICKEN-ORGAN	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55015LA	CHICKEN-ORGAN	25 COOKED-FRESH-FRIED	7F3476	P 0.020000	0.020000		100.00	0.020000
55015LA	CHICKEN-ORGAN	26 COOKED-FRESH-PICKLED,CORNEED,OR CURED	7F3476	P 0.020000	0.020000		100.00	0.020000
55015MA	CHICKEN-W/O SKIN	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55015MA	CHICKEN-W/O SKIN	22 COOKED-FRESH-BAKED	7F3476	P 0.020000	0.020000		100.00	0.020000
55015MA	CHICKEN-W/O SKIN	25 COOKED-FRESH-FRIED	7F3476	P 0.020000	0.020000		100.00	0.020000
55015MA	CHICKEN-W/O SKIN	31 COOKED-FRESH OR CANNED	7F3476	P 0.020000	0.020000		100.00	0.020000
55015MA	CHICKEN-W/O SKIN	53 COOKED-CANNED-BOILED	7F3476	P 0.020000	0.020000		100.00	0.020000
55015MB	CHICKEN+SKIN	21 COOKED-NFS	7F3476	P 0.020000	0.020000		100.00	0.020000
55015MB	CHICKEN+SKIN	25 COOKED-FRESH-FRIED	7F3476	P 0.020000	0.020000		100.00	0.020000



13544

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Chemical: Myclobutanil

PC Code:
128857

HED File Code: 11100 Other Chemistry Documents

Memo Date: 5/4/1998

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